PATENT SPECIFICATION

NO DRAWINGS

Inventors: RICHARD MOATS SHEELEY and GEORGE RODGER ALLEN, IR.

1.164.360



Date of Application and filing Complete Specification: 30 Nov., 1967. No. 54652167.

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ERRATUM

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Anns invention relates to a method of preparing substituted 11-aminodibenz-[b,f][1,4]oxazepines or thiazepines and substituted 11-aminodibenz[b,f][1,4]diazepines. The products of the method are useful for their (CNS) activity as tranquilizers and anti-depressants as well as analgesics.

The diszepines, oxazepines and thiszepines which may be prepared by the method of this invention may be represented by the following formula:

10

15

wherein R. and R. are the same or distinct and each is hydrogen, (C,—C,) alkyl, (C,—C,) alkyr, nitra, habren or Burromythyl: R. is hydrogen or (C,—C, alkyl), (C,—C) alkyland, (C,—C, alkyland,).

 $(C_i - C_e \ alkyl \ or \ \text{--}(hydroxy) \ (C_i - C \ \ alkyl: \ or \ \text{--}N$ when taken together is

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International Classification: -C 07 d 57/60; 87/54; 93/42; 99/02

COMPLETE SPECIFICATION

A process for Preparing Tricyclic Organic Compounds

We, AMERICAN CYANAMID COMPANY, a corporation organised and existing under the laws of the State of Matine, United States of America, of Berdan Avenue, Township of Wayne, State of New Jersey, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by

the following statement:

This invention relates to a method of preparing substituted 11-aminodibenz[b,f][1,4]oazazepines or thiazepines and substituted 11-aminodibenz-[b,f][1,4]diazepines. The products of the method are useful for their (CNS) activity as tranquilizer-

and anti-depressants as well as analogesics.

The diazepines, saxepines and thiazepines which may be prepared by the method of this invention may be represented by the following formula:



15

5



(T)

wherein R₁ and R₂ are the same or different and each is hydrogen, (C_1-C_6) alkyl, (C_1-C_6) alkoyy, nitro, halogen or trifluoromethyl; R₂ is hydrogen or (C_1-C_6) alkyl, (C_2-C_6) alkyl, (C_2-C_6) alkyl, (C_2-C_6) alkyl, (C_3-C_6) alkyl,

 $(C_r - C_\epsilon \text{ alkyl or } \circ \cdot \text{(hydroxy)} \ (C_r - C_\alpha) \text{ alkyl; or } - N \qquad \text{when taken together is}$

[Price

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4-(C,—C_s) alkyl)-1-piperazinyl, 4-[hydroxy-(C,—C_s) alkyl]-1-piperazinyl, 4-[dialkylamino-(C,—C_s) alkyl]-1-piperazinyl, piperdino, morpholino or 2,2-polymethy-lenchydrazino; and Z is oxygen, sulfur or >N-(C,—C_s) alkyl.

The compounds with which this invention is concerned are, in general, white

compounds with winds one increased in the control of the control o

and in aqueous solutions for parenteral injection.

The method of this invention involves the cyclization of substituted thioureas (II) to produce compounds (I) as illustrated by the following reaction scheme:

20 wherein R., R., R., R., and Z are defined as described hereinabove. The conversion of compounds II into compounds I is achieved by the use of phosphorus coxychoride, optionally, and preferably, in conjunction with phosphorus permoxide. An excess of the oxychloride may also serve as a suitable solvent. The cyclization reaction is generally carried out at temperatures of from 50°C to 150°C, with the preferred temperature being from 80°C to 110°C. The addition of other solvents which are inert under the reaction conditions may also be useful, such as beaucae, toluene and xylene. When the cyclization has been achieved, usually after heating from 30 minutes to 24 hours, the products (I) may be recovered by treating the reaction solution with water, followed by purification of the crude product by methods well known to the skilled in the attention.

to those skilled in the art.

The thioureas (II) which serve as the starting materials for (I) in the method of this invention may be prepared from substituted diphenyl ethers, substituted diphenyl suffices, and substituted diphenyl suffices, and substituted diphenylamines as set forth in the following reaction scheme:

wherein R_{12} R_{23} R_{23} R_{4} and Z are defined as hereinabove. In accordance with this reaction scheme, acylation of the substituted diphenyl ethers, substituted diphenyl sulfides and substituted diphenylamines with phenoxy thiocarbonyl chloride produces the thiocarbonilates (IV). Treatment of the latter compounds with an appropriate mono or diamine gives the thiomeas (II).

	The compounds prepared by the method of the present invention are physiologi-	
	cally active on the central nervous system. They show high activity as tranquilizers at	
	non-toxic doses and in some instances anti-depressant properties at dosage levels	
5	which produce neither overt stimulation nor depression.	
,	A useful test for tranquilizer activity consists of measuring the reduction of	5
	spontaneous motor activity in animals by means of an actophotometer (a photoelectric device for quantitatively measuring locomotor activity). Graded doses of the active	
	compounds prepared by the process of this invention are administered to groups of	
	mice, and the effective dosage range for a significant reduction of motor activity (a	
10	measure of transmitization; compared to control groups is aetablished. The use of	10
	reduced moter activity as a measure of tranquilizing activity has been described by W. D. Gray, A. C. Osterberg and C. E. Rauh, Archives Internationales et de	10
	W. D. Gray, A. C. Osterberg and C. E. Rauh, Archives Internationales et de	
	Incrapic, vor. 134, p. 196 (1901) and W. I. Kinnard and C. I. Carr. Inurnal of	
15	Pharmacology and Experimental Therapeutics, Vol. 121, p. 354 (1957). The anti-depressant properties of the compounds prepared by the method of the	
	present invention are evident by measuring their ability to counteract a depression	15
	induced in animals by the administration of tetrahenazine heyamate. Graded doses	
	of the active compounds of this invention are administered to groups of mice, and	
20	this is followed by administering a dose of tetrabenazine which is known to markedly depress the exploratory behaviour of normal mice. The anti-depressant treated	
20	depress the exploratory behaviour of normal mice. The anti-depressant treated	20
	treated with an ineffective anti-depressant agent, do not show this normal exploratory	
	behaviour, but show well known profound depression induced by tetrabenazine. The results from several dose levels are used to establish effective dose ranges. The	
25	and-depressant compounds prepared by the process of this invention show their de-	25
	SITADIC PRODUCTIES BY this procedure at dose levels which produce little or no un-	20
	toward reactions, such as ataxia or reduced spontaneous motor activity	
	In addition, some of the compounds prepared by the method of this invention	
30	show other valuable pharmaccutical properties, such as analgesic activity. The following examples describe in detail the preparation of representative sub-	
	stituted 11-aminodibenz,b,f][1,4]-oxazepines and thiazepines and substituted 11-	30
	aminodibenz[b,f][1,4]diazepines by the method of the present invention.	
	-	
	EXAMPLE 1	
35	Preparation of 11-(1-Piperidinyl)dibenz[b,f][1,4]oxazepine	
35	The compound, phenyl o-phenoxythiocarbanilate, and piperidine are heated in ethanol at reflux temperature, followed by removal of the solvent and recrystalliza-	35
	tion of from dilute ethanol. The product obtained is 2'-phenoxypiperidinethiocarbox-	
	aniide, melting point 135—136°C.	
40	2'-phenoxypiperidinethiocarboxanilide (1.0 g.) is refluxed with phosphorus pent-	
40	oxide (approx. 1 g.) in phosphorus oxychloride (5 ml.) for two hours. After cooling to	40
	room temperature, the reaction mixture is diluted with ether. The resulting precipi-	
	tate is washed several times with other, treated with aqueous potassium carbonate	
	solution (preferably with cooling) and the mixture extracted three times with ether. The ether solution is dried over anhydrous potassium carbonate, and upon evapora-	
45	uon of the solvent, yields a vellow solid which is recrystallized from bentane to give	45
	white crystals, melting point 98-100°C.	
	Example 2	
	Preparation of 11-Aminodibenz[b,f][1,4]oxazepine Using the procedure described in Example 1, and treating phenyl o-phenoxy-	
50	unocarpanilate with ammoniscal effer produces the product 1_/2_phenovyl_2_phenovyl	50
	unloured, meiling point 125-127°(after recrustallization from dilute mothers!	50
	rieating 1-(2-phenoxy)-phenylthionrea, prepared shows with phosphorus page	
	oxide in phosphorus oxychloride, gives the product as crystals, melting point 198—200°C.	
55	200°C.	
	Example 3	55
	Preparation of 11-(n-Butylamino)dihenz[h f][14]ovavenine	
	4-Aminogiphenyl ether hydrochloride (16.66 g 0.075 mole) is converted the	
AD.		
	solution is treated with a solution of 6.45 g. (0.0375 mole) of phenoxythiocarbonyl chloride in ether. The resulting solution is magnetically stirred at ambient tem-	60
	perature for about 21 hours; filtration gives white crystals of 2-aminodiphenyl ether	
	- Since Capacity of 2-annitotriplienty ether	

4	1,164,360	
5	hydrochloride. The solvent is removed from the filtrate to give an amber oil that crystallizes from hexane to give 12.4 g, of solid, melting point 95—98°C. A solution of 3.21 g, (10 moles) of phenyl o-phenoxylthicarbantlate prepared above, and 1.46 g, (20 moles) of n-bavylamine in 50 ml. of ethanol is heated at re-flux temperature for 50 minutes. The solvent is removed, and the residue crystallizes on trituration with hexane. Recrystallization of the product 1-bavyl-5-2-c phenoxylpenylyltionura from ether-hexane gives white crystals, melting point 80—	5
10	In the manner described in Example 1, treatment of 1-butyl-3-(2-phenoxy-phenyl)phiourea prepared above, with phosphorus pentoxide in phosphorus oxy-chloride gives crystals, melting point 67—71°C, after recrystallization from heptane.	10
15	Preparation of 11-(4-Methyl-1-pipenziny)dibenz[b,f] [1,4] oxazepine In the manner described in Example 1, treatment of phenyl o-phenoxythio- carbanilate with a molar equivalent of 1-methyl-pipenzine gives the product, 4- methyl-2-piencyl-pipenzin-indocarboxalilok, which forms white crystals, melting point 139—142°C., from dilute ethanol. Using the procedure described in Example 1, and treating 4-methyl-2-plenoxy- control of the procedure described in Example 1, and treating 4-methyl-2-plenoxy- control of the procedure described in Example 1.	15
20	v. Sing use procusation of procusing the procusing the procusing the procusing the procusing the procusing the procusing point 96—97°C, after recrystallization from heptane.	20
25	Preparation of 2-Chloro 11-[44-(β-Hydroxychyl)-1-piperaziny] dibenz [b,1][1,4]osazzine Dihydroxhoride Using the procedure described in Example 1 and treating phenyl 2-(p-chloro- phenoxythiocarbanilate with 1-(β-hydroxychyl)-piperazine gives the product 2-(p- chlorophenoxy-4-(2''n-ydroxychyl)-1-piperazinehlocartoxanilde, melting point	25
30	153—153°C, after recrystallization from ethyl acetate. Heating 2c · (p · chlorophenoxy) + 4 · (2c ·) - hydroxyethyl) - 1 - piperazine - thiocarboxanilide, prepared above, with phospherus pentoxide in phosphorus oxy-chloride gives the product which is dissolved in ether and treated with hydrogen chloride to furnish 2-chloro 11 · [4 · (β - hydroxyethyl) - 1 - piperazinyl] - thence (14 · (β - hydroxyethyl) - 1 - piperazinyl] - thence (14 · (β - hydroxyethyl) - 1 - piperazinyl] - thence (14 · (β - hydroxyethyl) - 1 - piperazinyl] - thence (15 · (β - hydroxyethyl) - 1 - piperazinyl] - thence (15 · (β - hydroxyethyl) - 1 - piperazinyl) - thence (15 · (β - hydroxyethyl) - 1 - piperazinyl) - thence (15 · (β - hydroxyethyl) - 1 - piperazinyl) - thence (15 · (β - hydroxyethyl) - 1 - piperazine (15 · (β - hydroxyethyl) - piperazine (15 · (β - hydroxyethyl) - 1 - piperazine (15 · (β - hydroxyethyl) - piperazine (15 · (β - hydroxyethyl) - piperazine (15 · (β - hydroxyethyl) - piperazine (15	30
35	EXAMPLE 6	35
40	Preparation of 2-Chloro-11-(4-methyl-1-pipenziaryldibenz [b.f]]1.4/onzapine Using the procedure described in Example 1, and treating phenyl 2-(p-chloro-phenoxy) thiocarbanilate with 1-methyl-pipenzine gives crystals of 2' · (p · chloro-phenoxy) - 4 · methyl - 1 · pipenziarchitocarboxanilide, melting point 145—147°C, after recrystallization from actone-hexame. When the procedure described in Example 1 is used and the starting material is 2' · (p · chlorophenoxy) - 4 · methyl - 1 · pipenziarchitocarboxanilide, the product obtained is buff-colored crystals, melting point 109—111°C.	40
45	Example 7 Preparation of 2-Chloro-11-(4-methyl-1-piperazinyl)dibenz	45
50	I the manuer described in Engage I, treatment of phenyl (p-chlorophenyl-thio)thiosenbullare with a under convolution of I-methylpieruzine gives the product. In the manuer described in Engage I, treatment of phenyl (p-chlorophenyl-thio)phenyl-thio)phenyl-thio)phenyl-thio)phenyl-thio)phenyl-thio)thio-thio-thio-thio-thio-thio-thio-thio-	50
55	Example 8 Preparation of 5-Methyl-11-(4-Methyl-1-piperazinyl)dibenz	55
60	In the manner described in Example 1, treatment of 2-amino-N-methyldi- phenylamine with phenoxythiocarbonyl chloride furnishes phenyl 2-(N-methylamlino)- thiocarboniates, which is obtained as an oil. In the manner described in Example 1, treatment of phenyl 2-(N-methylamlino)-	60

1,164,360 5

	thiocarbanilate, prepared above, with 1-methylpiperazine gives crystals, of 4-methyl- 2'-(N-methylanilino)-1-piperazinethiocarboxanilide, melting point 131—132°C., after recrystallization from heptane-ethyl acetate.	
5	Using the procedure described in Example 1, treatment of 4 - methyl - 2' - N - methylanilino - 1 - piperazinethiocarboxanilide with phosphorus pentoxide in phosphorus oxychloride gives crystals having melting point 119—120°C. after recrystallization from heptane.	5
	Example 9	
10	Preparation of 11-Morpholino-dibenz[b,f][1,4]thiazepine Hydrochloride In the manner described in Example 1, treatment of 2-aminodiphenyl sulfide with phenoxythiocarbonyl chloride furnishes white crystals, melting point 86—88°C, after recrystallization from ethanol.	10
15	Using the procedure described in Example 1, treatment of phenyl 2-(phenylthio)-caroanilate with morpholine in propanol gives crystals of 2° - (phenylthio) - 4 - morpholinehioarboxanilide, melting point 83—84°C, after recrystallization from ethanol-heptane.	15
20	A solution of 300 mg, of 2°Cphenythio)-4-morpholinethiocarboxanilide in 2ml of phosphorus oxychloride containing a drop of dimetrylfornamide is heated at reliux temperature for 5 hours. The product is isolated in the manner described in Example 1 and is obtained as cream-colored crystals, melting point 250—260°C, after recrystallization from ethyl acetatemethanol.	20
	EXAMPLE 10 Preparation of 11-(2-Dimethylaminoethylamino)dibenz[b,f][1,4]oxazepine	
25	Following the procedure described in Example 1 and treating phenyl ophenoxythiocarbanilate with unsym-dimethylethylene diamine in ethanol gives the product, 1 - (2 - dimethylaminoethyl) - 3 - (2 - phenoxy - phenyl)thiourea as a coloriess oil.	25.
30	Heating 1 - (2 - dimethylaminoethyl) - 3 - (2 - phenoxy - phenylthiourea with phosphorus oxychloride in the presence of dimethylformamide at refluxing temperatures produces 11 - (2 - dimethylaminoethylamino)dibenz[b,f][1,4]oxazepine, melting point 82—84°C.	30
	Example 11	
35	Preparation of 2-Chloro-11-[(3-dimethylaminocthylmethylamino]-dibenz [b,f][1,4] thiazepine Hydrochloride Using the procedure described in Example 1 and treating 2 - (p - chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with trimethyl ethylenediamine gives the product 1 - [2 - (p - the chlorophenyl - thio)thiocarbanilate with thio thiological with thiological with thiological with thiological with the product 1 - [2 - (p - thiological with thio	35
	chlorophenylthio)phenyl] - 3 - methyl - 3 - (2 - dimethylaminoethyl)thiourea, which is isolated as the hydrochloride salt, melting point 167—168°C.	
40	Heating 1 · [2 · (9 · chlorophemythio)phemy] · 3 methyl · 3 · (2 · dimethylaminocthyl) thioures hydrochloride, prepared above, with phosphorus pentoxide in plusphorus oxychloride gives the produce which is dissolved in ether and treated with hydrogen chloride to frumsh 2 · chloro · 11 · [(3 · dimethylaminocthyl) methylamino] · dibenz[bx][1,14] thiazzpine hydrochloride, methig point 196—197°C.	40
45	Example 12	
45	Preparation of 2-Chloro-11-[4-(3-dimethylaminopropyl)-1-piperazinyl] dibenz[b,f][1,4]osazepine In the manner described in Example 1, treatment of phenyl 2-(p-chlorophenoxy)-	45
* 0	thiocarbanilate with 1-(3-dimethylaminopropyl)piperazine gives an oil that is converted into the dihydrochloride upon treatment with ethereal hydrogen chloride.	
50	Crystallization from methanol gives $2'-(p$ - chlorophenoxy) - 4 - $(3$ - dimethyl - aminopropyl) - 1 - piperazinethiocarboxanilide dihydrochloride, melting point 208—210°C.	50
55	Following the process of Example 1, and heating 2^{\prime} – $(p$ - chlorophenoxy) - 4 – (3 - dimethylaminopropy) - 1 – piperazinelinearboxaniide dihydrochloride, prepared above, with phosphorus pentoxide in phosphorus oxychloride, the product 2 - chloro - 11 - $[4$ – (3 - dimethylaminopropyl) - 1 – piperazinyl]dibena[b,i][1,4] oxazephe is obtained.	55
	Example 13	
60	Preparation of 2-Chloro-11-(4-morpholinyl)dibenz[b,f][1,4]thiazepine Using the procedure described in Example 1 and treating phenyl 2-(b-chloro-	60

5	phenythio/thiocarbanilites with morpholine gives the product 2°-(p-chlorophenyl- thio)-4-mepholine-thiocarboanilitie, melting point 63—67°C. Heating 2°- (p-chlorophenylithio)-4 - morpholinethicarboanilide, prepared above, with phosphorus pentoside in phorphorus oxychloride gives the product 2- chloro-11-(4-morpholinyl)dibenz[b,f][1,4]thiazepine as crystals, melting point 148— 150°C.	5
10	EXAMPLE 14 Preparation of 2-Chloro-11-(1-piperdimylamno)dibenz[bf][1,4]diazepine Following the procedure described in Example 1 and treating phenyl 2-(p-chlorophenylthio)chlocarbanilate with N-aminopiperdine furnishes the product 1-(2-p-chlorophenylthio)-phenyl -3-piperdino-thiourea, melting point 1835—160°C, after recrystalization from ethanol.	10
15	D83—10 ¹ C ₂ after (Lysalization for Charlette (Lysalization for Charlette) = 3 - piperidino - thioures, pre- pared above, with phosphorus pentoxide in phosphorus oxychloride gives the pro- duct 2 - chloro - 11 · (1 - piperidinylamino)dibenz[b,f][1,4]thiazepine as crystals, melting point 153—154°C.	15
20	HEXAMEL 15 Preparation or 11-(β -Hydroxyethylamino)tilbenz[b,f] [1,4]ocazepine Using the procedure described in Example 1 and treating phenyl 2-phenoxythio- carbanilate with echanolamine gives the product 1 - (β - hydroxyethyl) - 3 - (Z' - phenoxy -phenylythourea.	20
25	pacmosy - pacity/innourae, prepared Henting 1 - (\hat{g} - hydroxyethyl) - 3 - ($2'$ - phenoxy - phenyl)thiourae, prepared above, with phosphorus pentoxide in phosphorus oxychloride gives the product 11-(\hat{g} -hydroxyethylamino)dibenz[b,f] [1,4]oszzepine as crystals, melting point 136—139°C.	25
30	Preparation of 11-(3-Dimetribaninepropy) dibenz [h.f] [1,4] oxazepine Following the precedure described in Example 1 and treating phenyl 2-penylthocarbaniles with c-dimethylaminopropylamine furniless the product 1-(3 - dimethylaminopropyl) - 3 - (2 - phenoxy - phenyl) dibourca, melting point 113—114-Co, after necrystallization from ethanol-heptane. Heating 1 - (3 - dimethylaminopropyl) - 3 - (2 - phenoxy - phenyl) thiourca, prepared above, with phosphorus pearoxide in phosphorus oxychloride gives the product as crysvalls, melting point 108—109-Co.	30
35	EXAMPLE 17 Preparation of 11-Allylaminodiberz[b,f] [1,4]oxazepine Hydrochloride Using the procedure described in Example 1 and treating phenyl 2-phenoxythio- carbanilate with allylamine gives the product 1 - allyl - 3 - (2 - phenoxyphenyl) -	35
40	theoures as an oil. Heating 1 - allyl - 3 - (2 - phenoxyphenyl) - thioures, prepared above, with phosphorus pentexide in phosphorus oxychioride gives the product I1-allylaminodibenz[bhf]1.4]oxzepine, the hydrochloride of which is obtained as crystals, melting point 20°C, with decomposition.	40
45	EXAMPLE 18 Preparation of 2-Fluoro-11-(4-methyl-1-piperazinyl)dibenz[b,f][1,4] oxazepine	45
50	Following the procedure described in Example 1, and treating phenyl 2 - $(\rho - \mu \cos \rho \tan \phi)$ induced phenoxyl-inductabuliste with 1-melhylingenzaine furnishes the product $2' - (\rho - \mu \cos \phi)$ fluorophenoxy) - 4 - methyl - 1 - piperazinethiocarboxanilide. Heating $2' - (\rho - \mu \sin \phi)$ fluorophenoxy) - 4 - methyl - 1 - piperazinethiocarboxanilide, prepared above, with phosphorus pentoxide in phosphorus oxychloride gives the product as crystals, metiting point $2^4 - 4^4 - 8^4 - $	50
55	EXAMPLE 19 Preparation of 2-Bromen 1:44-methyl-1-piperazinyl)dibenz [16,1] 1.44-methyl-1-piperazinyl)dibenz [16,1] 1.45 methyl-1-piperazinyl)dibenz Using the procedure described in Example 1 and treating phenyl 2 - (p - bromophenoxy) - 4+ methyl - 1- piperazine-pives the product 2' - (p - bromophenoxy) - 4+ methyl - 1 - piperazine-thiocarboxa-dilike. Heating 2' - (p - bromophenoxy) - 4 - methyl - 1 - piperazine-thiocarbox -	55

	anilide, prepared above, with phosphorus pentoxide in phosphorus oxychloride gives the product as crystals, melting point $95-99^{\circ}C$.	
	Example 20	
5	Preparation of 2-Methyl-11-(4-methyl-1-piperazinyl)dibenz [bf][1.4]oxazepine Following the procedure described in Example 1 and treating above 2 - (6.	5
10	tolyloxy/thiocuthenilate with 1-methylipicrazine furnishes the product 2' - (p - tolyloxy) - 4 - methyl - 1 - piperazinethiocurbosanilide. Heating 2' - (p - tolyloxy) - 4 - methyl - 1 - piperazinethiocurboxanilide, prepared above, with phosphorus pentoxide in phosphorus oxychloride gives the product as crystals, melting point 130—131°C.	10
	Example 21	
15	Preparation of 8-Chloro-11-(4-methyl-1-piperazinyl)dibenz [bsf] [1.4]oxazepine Using the procedure described in Example 1 and treating phenyl 5 - chloro - 2 - phenoxythiographenilete with 1 methylicing and the control of the contro	15
20	2' - phenoxy - 4 - methyl - 1 - piperazinethiocraborallide. Heating 5' - chloro - 2' - phenoxy - 4 - methyl - 1 - piperazinethiocraboxanilide, prepared above, with phosphorus pentoxide in phosphorus oxychloride gives the product as crystals, melting point 165—166°C.	20
	Example 22	
25	Preparation of 6-Chiloro-II-(4-methyl-1-piperazinyl)dibenz [by][I-(4)nexpip-iperazinyl)dibenz [by][I-(4)nexpip-iperazinyl)dibenz [by][I-(4)nexpip-iperazinezinezinezinezinezinezinezinezinezine	25
30	EXAMPLE 23	20
	Preparation of 4-Chloro-11-(4-methyl-1-piperazinyl)dihenz	30
35	Using the procedure described in Example 1 and treating phenyl 2 - (o - chlorophenoxy)-thiocarbanilate with 1-methylpiperazine gives the product 2' - (o - chlorophenoxy) - 4 - methyl - 1 - piperazinethiocarboxanilide. Heating 2' - (o - chlorophenoxy) - 4 - methyl - 1 - piperazinethiocarboxanilide, product acrystals, melting point 173—174°C.	35
	Example 24	
40	Preparation of 11-(4-Methyl-1-pipenzinyl)-8-trifluore-methyldibenz [Js.][1]-(4-Methyl-1-pipenzinyl)-8-trifluore-methyldibenz [Js.][1]-(4-Methyl-1-pipenzinyl-1-pipenzinyl-1-pipenzinyl-1-pipenzinyl-1-pipenzinyl-1-pipenzinthio-ductiv-1-pipenzint	40
45	auct y - trittuoromethyl - 2' - (phenylthio) - 4 - methyl - 1 - piperazinethio - carboxanilide, Heating 5' - tritluoromethyl - 2' - phenylthio - 4 - methyl - 1 - piperazine - thiocarboxanilide, prepared above, with phosphorus pentoxide in phosphorus oxy-	45
50	thiocarboxanlide, prepared above, with phosphorus pentoxide in phosphorus oxy- chloride gives the product which is dissolved in ether and treated with hydrogen chloride to furnish 11 - (4 - methyl - 1 - piperazinyl) - 8 - trifluoromethyl - diberz[b,f][1,4]thiazepine dihydrochloride as crystals, melting point 192°C, with de- composition.	50
55	EXAMDE 25 Preparation of 2-Methoxy-11-4-methyl-1-piperidinyl)dibenz [16-f] [1-4]thhazpine Using the procedure described in Example 1 and treating phenyl 2-(ρ -methoxy-phenylthio)dibicarbanilate with piperidine gives the product $2' \cdot (\rho$ - methoxy-phenyl-thio) - 1 - piperidinenthiox-box-method (Heating 2' - (ρ - methoxy-phenylthio) - 1 - piperidinenthiocarboxanilide, pre-	55

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hexane.

pared above, with phosphorus pentoxide in phosphorus oxychloride gives the product as crystals, melting point 116-117°C.

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EXAMPLE 26 Preparation of 2-Nitro-11-(4-methyl-1-piperazinyl)dibenz [b,f][1,4]oxazepine

1 - Methyl - 4 - piperazimenhocatowyl chloride, prepared from I-methylpiperazine and thiophsogene is treated with o-aminopherol to give I - methyl-4 - (o - hydroxyphenythhocatomory)piperazine. This preduct is treated with 4chloronitrobenzene in dimethylaceratine in the presence of potassium cathonate. A Methyl - 2' - (o - nitrophenoy)piperazine thiocathocanildie is thereby obtained.

Methyl - 2' - (p' - mitorpiends); preparation to the property of the property

WHAT WE CLAIM IS:—

1. A method of preparing a compound of the formula:

wherein R_1 and R_2 are the same or different and each is hydrogen, (C_1-C_4) alkyl, (C_1-C_4) alkoy, nitro, halogen or trifluoromethyl; R_2 is hydrogen or (C_1-C_2) alkyl; R_4 is hydrogen, (C_1-C_4) alkyl, (C_2-C_4) alkenyl, (C_1-C_4) alkylamino (C_1-C_5)

alkyl, or
$$\omega$$
-(hydroxy(C₁—C₀) alkyl or —N when taken together is 4-[(C₁—C₀)

alkyl]-1-piperazinyl, 4-[hydroxy-(C,—C,alkyl]-1-piperazinyl, 4-[dialkylamino (C,—C,alkyl]-1-piperazinyl, piperidino, morpholino, 2,2-polymethylenchydrazino; and Z is oxygen, sulfur or >N-(C,—C,) alkyl, the method comprising heating a thiourea of the formula:

$$R_3$$
 R_4
 R_4
 R_1
 R_2

wherein R₁, R₂, R₃, R₄ and Z are as defined above, with phosphorus oxychloride, optionally in conjunction with phosphorus pentoxide.

2. A method according to Claim 1, wherein the thiourea is 4 - methyl - 2' - phenoxypiperazinethiocarboxamilide and the product obtained is 11 - (4 - methyl - meth

1 – piperazimyl-Jólhenz [h.][1.4] osazepine.
3. A method according to Claim 1, wherein the thiourea is 2' – (p – chloro-phenoxy) 4 – methyl 1 – piperazinethiocarboxanilide and the product obtained is 2 – chloro – 11 – (4 – methyl – 1 – piperazinyl-bibezz [b.][1]. Joszazepine.
4. A method according to Cleim 1, wherein the thiourea is 4 – methyl – 2' –

4. A method according to Calim 1, whetein the influence 1 and the product obtained is 2 - chloro-plenylthio) piperazine-fuccioro-planel in 1 - (4 - methyl - 1 - piperazinyl(dibenz[6,f][1,4] lthazepine.

5. A method according to Claim 1, wherein the thiourea is 1 - (2 - dimethyl - aminochiyl) - 3 - (2 - phenoxy - phenyl) historica and the product obtained is 11 - 40 (2 - dimethylaminochyl)dibenz [b.f.] [1,4] (oxazepine.

1.164.360

6. A method according to Claim 1, substantially as described in any one of the Examples herein.

7. A substituted diazepine, oxazepine or thiazepine whenever prepared by a

Process according to any preceding claim.

8. A pharmaceutical preparation comprising a compound according to Claim 7 together with a pharmaceutically acceptable carrier or diluent.

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